

OFFICE OF NAVAL RESEARCH Contract[N60014-77-C-6231, Task No. NR 053-640 TECHNICAL REPERT NO. 9 Crystal Structure and Properties of Tetrathiafulvalenium Triiodide. by
Robert C. /Teitelbaum, la, b, e Tobin J. /Marks la, 2 and
Carroll K. Johnson Id
Prepared for Publication
in
The Journal of the American Chemical Society
Northwestern University Department of Chemistry Evanston, Illinois 60201 Evanston
3 December 3, 1979 730 C

Reproduction in whole or in part is permitted for any purpose of the United States Government

This document has been approved for public release and sale; its distribution is unlimited

This statement should also appear in Item 10 of Document Control Data - DD Form 1473. Copies of form available from cognizant contract administrator.

26\$8\$5 **80 2 2**5

unclassified

SECURITY CLASSIFICATION OF THIS PAGE (When Date Entered)

REPORT DOCUMENTATION PAGE	BEFORE COMPLETING FORM
Technical Report No. 9	D. 3. RECIPIENT'S CATALOG NUMBER
4. TITLE (and Subilile)	5. TYPE OF REPORT & PERIOD COVERED
Crystal Structure and Properties of Tetrathiafulvalenium Trijodide	Interim, 1979
1 ett autatutvatenium 11 noutue	6. PERFORMING ORG. REPORT NUMBER
7. AUTHOR(a)	NO0014-77-C-0231
9. PERFORMING ORGANIZATION NAME AND ADDRESS	10. PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS NR-053-640
11. CONTROLLING OFFICE NAME AND ADDRESS	December 3, 1979
	13. NUMBER OF PAGES
14., MONITORING AGENCY NAME & ADDRESS(II different from Controlling Office)	15. SECURITY CLASS. (of this report)
	unclassified
	15. DECLASSIFICATION/DOWNGRADING SCHEDULE
17. DISTRIBUTION STATEMENT (of the abetract entered in Block 20, if different in	om Rapóri)
18. SUPPLEMENTARY NOTES	
19. KEY WORDS (Continue on reverse side if necessary and identify by block number)	
Polyiodide TTF	
Conductive material Organic conduct	tor
Mixed valence material	
Tetrathiafulvalene	
Slow cooling of a hot acetonitrile solution of te iodine yields a mixture of crystalline products inc of high iodine composition: TTF·I;. A single crystals been carried out on these crystals of symmetric crystals.	luding monoclinic crystals stal X-ray structure analysis

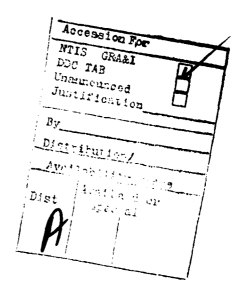
DD 1 JAN 73 1473 EDITION OF 1 NOV 65 IS OBSOLETE S/N 0102-014-6601 |

unclassified

SECURITY CLASSIFICATION OF THIS PAGE (When Date Entered)

LLUMITY CLASSIFICATION OF THIS PAGE(When Data Entered)

of triiodide ions. Consistent with the structural results, single crystal measurements indicate that this material exhibits high electrical resistivity with $\sigma_{\rm RT}$ < 9 x 10⁻⁹ Ω^{-1} cm⁻¹.



unclassified

SECURITY CLASSIFICATION OF THIS PAGE(When Date Entered)

4800000

By acceptance of this article, the publisher or recipient acknowledges the U.S. Government's right to retain a nonexclusive, royalty-free license in and to any copyright covering the article.

Contribution from the Department of Chemistry and the Materials Research Center
Northwestern University
Evanston, Illinois 60201 and the Chemistry Division
Oak Ridge National Laboratory
Oak Ridge, Tennessee 37830

CRYSTAL STRUCTURE AND PROPERTIES OF TETRATHIAFULVALENIUM TRIIODIDE

by Robert C. Teitelbaum, *Tobin J. Marks*la, 2 and Carroll K. Johnson* 1d

ABSTRACT

Slow cooling of a hot acetonitrile solution of tetrathiafulvalene (TTF) and iodine yields a mixture of crystalline products including monoclinic crystals of high iodine composition: TTF· Is. A single crystal X-ray structure analysis has been carried out on these crystals of symmetry P^2 in with 4 molecules per unit cell. The cell parameters are $a = 9.411(2) \, \text{Å}$, $b = 18.787(3) \, \text{Å}$, $c = 7.771(1) \, \text{Å}$, $\beta = 103.2(1)$, and $V = 1374(1) \, \text{Å}^3$. The structure consists of integrated stacks along the

02 pul 1

<102> axis with TTF⁺ dimers interspersed between pairs of tri-iodide ions. Consistent with the structural results, single crystal measurements indicate that this material exhibits high electrical resistivity with $\sigma_{\rm RT}^{-2}$ < 9 x 10⁻⁹ Ω^{-1} cm⁻¹.

INTRODUCTION

Considerable attention in recent years has been focused on the chemistry and physics of materials which exhibit highly anisotropic electrical, optical, and magnetic properties. Of particular interest have been organic and metal-organic solids which exhibit metal-like charge transport along one crystalline axis. The discovery of the "organic metal" tetrathiafulvalene-tetracyanoquinodimethane, TTF-TCNQ, prompted a number of studies of TTF as an electron donor and precursor for other highly conductive solids. Halogen oxidation, for example, yields a variety of charge transfer salts.

One direction of activity in our Laboratories and others has involved the iodine-containing TTF salts, TTF· I_x . The stoichiometries prepared to date range from x=0.71 to 3.0, but detailed single crystal characterization has been restricted to a limited number of low iodine phases. The most conductive of these is TTF· $I_{q,71}$ with a room temperature conductivity of ca. 300 Ω^{-1} cm⁻¹ along the needle axis. This material possesses an ordered, modulated crystal structure containing stacks of eclipsed, partially oxidized TTF molecules, and parallel chains of I^- ions. Two additional phases of intermediate stoichiometry with x ~2.0 have been isolated which vary only in the degree of order-disorder. The first TTF· I_2 structure (space group Immm) contains disordered chains of triiodide ions and disordered TTF stacks $I(TTF^{+0.7})(I_3^-)_{0,7}I_3^{-6}$, Preliminary X-ray crystallographic investigations sug-

gest that the second TTF. L phase(space group Fddd) is a superstructure of the first TTF. L phase with ordered TTF stacks, but again with disordered triiodide ions. Full expositions of the TTF. I2 structures, spectroscopy and transport properties are in preparation. In view of the accumulating knowledge on the above TTF-iodine complexes, it would clearly be of interest to investigate, for comparative purposes, the phase of greatest iodine content. This would provide information on how the crystal structure, 8 degree of partial oxidation, and transport properties adjust to the increased dopant level. Such information might also shed light on how large amounts of iodine metal glyoximate and are sometimes accomodated in phthalocyanine 10 lattices. Here, again, phases with very high iodine content have been isolated. 9, 10 We report here the synthesis, X-ray diffraction structure determination, and single crystal conductivity of the high iodine TTF phase, TTF · I₃.

EXPERIMENTAL

Synthesis of TTF· I_3 . Tetrathiafulvalene triiodide was prepared under a nitrogen atmosphere by adding dropwise a solution of 80 mg (0.39 mmole) TTF (Aldrich Chemical Co.) in 10 ml acetonitrile (freshly distilled from P_2O_5 under nitrogen), to a stirring solution of 120 mg. (0.47 mmole) triply sublimed I_2 in 10 ml of freshly distilled acetonitrile. Sufficient acetonitrile was added until the total volume was 50 ml . The solution was heated to 50° C with constant stirring until all of the solid present dissolved. Upon slow cooling to room temperature, a mixture of crystals of TTF· $I_{0,71}$, TTF· I_2 , and in trace amounts, TTF· I_3 resulted. The product was then washed with freshly distilled benzene and dried with a stream of nitrogen. The morphology of the TTF· I_3 crystals are sufficiently different from the other phases to be easily separated by inspection. Insufficient quantities of material were obtained for elemental analysis.

<u>X-ray Diffraction Study.</u> TTF·I₃ has 4 molecules per unit cell with space group symmetry $P2_1/n$. Least-squares refinement of cell parameters using 10 high order reflections gave the unit cell dimension $\underline{a} = 9.41l(2)$, $\underline{b} = 18.787(3)$, and $\underline{c} = 7.77l(1)$ Å, with $\beta = 103.2(1)^{\circ}$ and V = 1374(1)Å³.

Single crystal intensities from a polyhedral crystal were collected on the Oak Ridge computer-controlled diffractometer using filtered

molybdenum radiation. Of the 3915 unique reflections measured, 3284 had intensities greater than one sigma. A twin or perhaps an unidentified second phase was present and caused at least 75 reflections, mainly in the (hk0)zone, to have excessive observed intensity. The (0k0) reflections were all affected. An absorption correction applied to the data gave transmittancy factors of 0.13 to 0.17.

Structure Solution and Refinement. The three iodine atom positions were found using the direct-methods program MULTAN of Germain, Main and Woolfson. These matched the heavy-atom vectors of the Patterson map. The sulfur and carbon atoms were then found using a Fourier map computed with phases based on iodine atom positions. The four hydrogen atoms were placed in idealized positions and their positional parameters included in the refinement.

Refinement of all nonhydrogen atoms with anisotropic thermal motion and hydrogen atom positions led to an R index of 0.078 on F^2 for all 3915 reflections. The refinement was based on 3090 reflections with 75 reflections deleted because of the twinning probelm and reflections < 1.5 sigma omitted. The $R(F^2)$ index for the reflections used in the refinement was 0.054. Tables of the final $|F_0|$ and $|F_c|$ values for the 3915 reflections are available upon request.

Atomic scattering factors for S, C, H, and the central I were taken directly from the International Tables of X-ray Crystallography, Volume IV. Scattering factors for the two outer iodines in the tri-iodide were taken as the average of the scattering factors for I° and I¯, since it has been shown that the negative charge in the triiodide ion is distributed among the outer iodines, while the central iodine is approximately neutral. At The real and imaginary anomalous-dispersion scattering-factor correlation of Cromer and Liberman were included.

Electric Conductivity Measurements. The samples examined for do conductivity studies (using the apparatus described previously 9) were nearly hexagonal prisms, with all dimensions on the order of a few tenths of a millimeter. The small crystal size permitted only two probe conductivity measurements, though orientations both parallel and perpendicular to the hexagonal plane could be examined. Samples were mounted by pressure contact between two spring-loaded brass pins.

RESULTS AND DISCUSSION

The reaction of TTF with iodine in acetonitrile, followed by slow cooling, yields $\mathrm{TTF} \cdot \mathrm{I}_{\mathrm{X}}$ phases of various stoichiometries (see Experimental Section for methodology). The small quantities of $\mathrm{TTF} \cdot \mathrm{I}_{\mathrm{S}}$ crystals formed were mechanically separated from the other phases.

Description of the Structure. The final atomic coordinates and anisotropic temperature factors are given in Table I. A stereoscopic view of the unit cell contents is shown in Figure 1 and the Li ion close contacts are illustrated in Figure 2. The crystal structure consists of integrated stacks of cofacial TTF⁺ dimers in alternation with pairs of triiodide ions. The TTF cation is distorted somewhat from planarity, with all sulfur atoms 0.01-0.03 Å below (i.e. toward the other molecule of the dimer) and all carbon atoms 0.0-0.03 Å above the least-squares best plane. The spacing between least-squares best planes for the TTF⁺ ions within the dimer is 3.40(1) Å, but the S-S distances of 3.351(2) and 3.373(2) Å are somewhat shorter. The molecular packing in TTF-I3 is reminiscent of that communicated for TTF. Br(and isomorphous TTF. Cl), also integral oxidation state salts. 4i, j, 16 In these materials, integrated stacks of (TTF)₂⁺² dimers and halide ions also occur. The interplanar spacing between members of the dimer appears to be somewhat larger in the present case (3.40(1) \mathring{A}) than in TTF-Br (3.34 \mathring{A}).

These interplanar spacings are considerably shorter than the TTF spacings of 3.55 (1) Å found in the segregated stack structures of TTF· $I_{0,71}$ and TTF· I_2 . The spacings in the present case are also shorter than in the eclipsed $(TTF)_2^{+2}$ dimers of $(TTF)_2Ni(S_4C_4H_4)$ (here the S-S distance is given as 3.48 Å). The direct axis fractional components of the TTF⁺ plane normal in TTF· I_3 are (0.0679, -0.0099, 0.1166) which is approximately along the <102> direction. The TTF⁺ ions are almost exactly superimposed with a slight lateral offset of about 0.25 Å perpendicular to the long axis of the molecule.

The molecular structural parameters for TTF·I₃ are summarized in Table II. In Figure 3, the bond distances and angles within the tetrathiafulvalenium ion are illustrated. These indicate that the cation is rather symmetrical, even though this is not required by the crystallographic symmetry. A comparison of the TTF⁺ structural data with that of other TTF cations as well as with neutral TTF is set out in Table III. It can be seen that oxidation is accompanied by lengthening of the exocyclic double bond (a) and shortening of the contacts between the two carbon atoms involved in the exocyclic double bond and the sulfur atoms (b). The metrical parameters for TTF⁺ in the present study are experimentally indistinguishable from those in TTF·Br. ¹⁶

The $TTF \cdot I_3$ crystal structure also contains parallel, zig-zagging chains of triiodide ions. The arrangement of the triiodide ions is illustrated in Figure 2. The triiodide ion deviates slightly from

linearity with an I-I-I angle of 175.40(2). This slight bending is frequently observed in structures of triiodides (Table IV) and reflects the rather flat potential energy surface for molecular distortion. The fact that the terminal iodine atoms of the L₃ ion are slightly displaced toward the positively charged (TTF)₂⁺² dimer can be explained on an electrostatic basis. The negative charge in L_3^- is localized principally on these two atoms. The slight asymmetry in the I-I distances, 2.908(1) and 2.953(1) Å, presumably also reflects the environment and ready deformability of the I_3 ion. The I-S contacts are illustrated in Figure 2, and the closest distances are in the range 3.636(2) - 3.989(2) Å. These values are typical of the I-S contacts found in TTF·I, systems. In the more closely packed TTF· $I_{0.7}$ structure, the I-S contacts are as small as 3.53 Å. As shown in Figure 2, the closest contact between proximate I₃ ions is 4.335(1) Å.

Electrical Conductivity. The room temperature dc conductivities of two single crystal samples of $TTF \cdot I_3$ were measured parallel to the hexagonal plane and found to have values of $6 \times 10^{-9} \ \Omega^{-1} \ cm^{-1}$ and $9 \times 10^{-10} \ \Omega^{-1} \ cm^{-1}$. The dc conductivity of the latter sample was also measured perpendicular to the hexagonal plane. Here the geometry put the precise value of the conductivity beyond the limita-

tions of the instrumentation, although an upper limit of 9 x 10^{-9} Ω^{-1} cm⁻¹ was determined. Finally, a small polycrystalline sample was examined and found to have $\sigma = 8 \times 10^{-10}$ Ω^{-1} cm⁻¹.

CONCLUSIONS

This work shows the high iodine phase of the TTF·I_ system x = 3, to be an integrated stack material with integral formal oxidation states, i.e. TTF⁺L₁. Not surprisingly, such a structure gives rise to high electrical resistivity. As in the case of the other known integral oxidation state tetrathiafulvalenium salts, 4h, i the structural motif in TTF·I₃ consists of integrated stacks of alternating $(TTF)_2^{+2}$ dimers and pairs of counter anions. Those factors which govern the stability of various possible crystal structures in TTF salts have been discussed at length. 4h, i, 19, 20 It appears that the present structural result can be best rationalized in terms of the electrostatic destabilization with respect to the partially oxidized situation of an integral oxidation state - (+1) segregated stack structure. 4h, i, 19 In addition, the close interplanar separation in the $(TTF)_2^{+2}$ suggests enhanced covalent bonding $^{20, 21}$ between the cationic moieties in the integral oxidation state case.

ACKNOWLEDGMENTS

The chemical and physicochemical research at Northwestern

University was generously supported by the Office of Naval Research,

and by the NSF-MRL program through the Materials Research Center of

Northwestern University (grant DMR76-80847). The crystallographic studies at Oak Ridge National Laboratory were generously supported by the Division of Materials Sciences, U.S. Department of Energy, under contract W-7405-eng-26 with the Union Carbide Corporation. We also thank Mr. M. Mc Clure for assistance with the conductivity measurements and Dr. S.J. La Placa of IBM Watson Laboratories for communicating his structural data on TTF·Br.

Supplementary Material Available: A listing of structure amplitude tables (2) pages). Ordering information is given on any current masthead page.

REFERENCES

- (1) a. Department of Chemistry and the Materials Research Center, Northwestern University.
 - b. Visiting Scientist, Oak Ridge National Laboratory.
 - c. Present address: Research Laboratories, Eastman Kodak Co., Kodak Park, Rochester, NY 14650.
 - d. Chemistry Division, Oak Ridge National Laboratory.
- (2) Camille and Henry Dreyfus Teacher-Scholar.
- (3) a. Devreese, J.T.; Evrard, V.E.; Van Doren, V.E., Highly Conducting One-Dimensional Solids, Plenum Press, N.Y., 1979.
 - b. Torrance, J.B. Accts. Chem. Res, 1979, 12, 79-86.
 - c. Miller, J.S.; Epstein, A.J., eds., Synthesis and Properties of Low-Dimensional Materials, Ann. N.Y. Acad. Sci., 1978, 313.
 - d. Keller, H.J., ed. Chemistry and Physics of One-Dimensional Metals, Plenum Press, New York, 1977.
 - e. Miller, J.S.; Epstein, A.J. <u>Prog. Inorg. Chem.</u>, <u>1976</u>, 20, 1-1 1-151.
 - f. Keller, H.J., ed. Low Dimensional Cooperative Phenomena, Plenum Press, N.Y. 1975.
 - g. Soos, Z.G.; Klein, D.J. in Molecular Associations, Foster, R., ed., Academic Press, N.Y., 1975, chapt. 1.
- (4) a. Warmack, R.J.; Callcott, T.A.; Watson, C.R. Phys. Rev. B. 1975, 12, 3336-3338.
 - b. Somoano, R.B.; Gupta, A.; Hadek, V.; Datta, T.; Jones, R.; Herman, A.M. J. Chem. Phys., 1975, 63, 4970-4976.
 - c. Johnson, C.K.; Watson, Jr., C.R.; Warmack, R.J. Abstr. Amer. Cryst. Assoc., 1975, 3, 19.

The same

- d. Daly, J.J.; Sanz, F. Acta. Crystallogr. Sec. B., 1975, 31, 620-
- e. LaPlaca, S.J.; Corfield; P.W.R.; Thomas, R.; Scott, B.A. Solid State Commun., 1975, 17, 635-638.
- f. LaPlaca, S.J. Bull. Amer. Phys. Soc., 1975, 20, 496.
- g. Johnson, C.K.; Watson, C.R., Jr.; Warmack, R.J. Chem. Div. Ann. Prog. Rep., Oak Ridge National Laboratory, 1976, 102-103.
- h Johnson, C.K.; Watson, C.R., Jr. J. Chem. Phys., 1976, 64, 2271-2286.
- i. Scott, B.A.; La Placa, S.J.; Torrance, J.B.; Silverman, B.D.; Welber, B. J. Amer. Chem. Soc., 1977, 99, 6631-6639.
- j. Scott, B.A.; LaPlaca, S.J.; Torrance, J.B.; Silverman, B.D.; Welber, B. in reference 3c, p. 369-376.
- (5) For 129 I Mossbauer studies see:
 - a. Marks, T.J. in reference 3c, p. 594-616.
 - b. Teitelbaum, R.C.; Marks, T.J. manuscript in preparation.
- (6) Marks, T.J.; Teitelbaum, R.C.; McClure, M.S.; Kannewurf, C.R.; Johnson, C.K.; Ruby, S.L.; Bull Amer. Phys. Soc., 1979, 24, 233.
- (7) a. Resonance Raman studies indicate that the major polyiodide constituent is a slightly distorted triiodide. 6, 7b
 - b. Teitelbaum, R.C.; Marks, T.J. unpublished results.
 - c. Johnson, C.K., manuscript in preparation.
- (8) For discussions of the structural chemistry of low-dimensional materials see:

- a. Kistenmacher, T.J. in reference 3c, p. 333-342.
- b. Megtert, S.; Pouget, J.P.; Comes, R. in reference 3c, p. 234-243.
- c. Stucky, G.D.; Schultz, A.J.; Williams, J. M. Ann. Rev. Mater. Sci., 1977, 7, 301-339.
- d. Bespalov, B. P.; Titov, V. V. Russ. Chem. Rev., 1975, 44, 1091-1108.
- e. Dahm, D.J.; Horn, P.; Johnson, G.R.; Miles, M.G.; Wilson, J.D. J. Cryst. Mol. Struct., 1975, 5, 27-34.
- f. Herbstein, F.H. Perspect. Struct. Chem., 1971, IV, 166-395.
- (9) a. Cowie, M.A.; Gleizes, A.; Grynkewich, G.W.; Kalina, D.W.; McClure, M.S.; Scaringe, R.P.; Teitelbaum, R.C.; Ruby, S.L.; Ibers, J.A.; Kannewurf, C.R.; Marks, T.J. J. Amer. Chem. Soc., 1979, 101, 2921-2936.
 - Brown, L.D.; Kalina, D.W.; McClure, M.S.; Ruby, S.L.;
 Schultz, S.; Ibers, J.A.; Kannewurf, C.R.; Marks, T.J.
 J. Amer. Chem. Soc., 1979, 101, 2937-2947.
- (10) a. Schramm, C J.; Stojakovic, D.R.; Hoffman, B.M.; Marks, T.J. Science, 1978, 200, 47-48.
 - b. Petersen, J.L.; Schramm, C.S.; Stojakovic, D.R.; Hoffman, B.M.; Marks, T.J. J. Amer. Chem. Soc., 1977, 99, 286-288.
 - c. Scaringe, R. P.; Schramm, C.J.; Stojakovic, D.R.; Hoffman, B.M.; Ibers, J.A.; Marks, T.J., submitted for publication.
 - d. Stojakovic, D.R. Ph.D. Thesis, Northwestern University, August 1977.
- (11) Germain, G.; Main, P.; Woolfson, M.M. Acta. Crystallogr., Sect. A, 1971, 27, 368-376.
- (12) See paragraph at end of paper regarding supplementary material.

· Alexander

- (13) Ibers, J.A.; Hamilton, W.C., eds. International Tables for X-ray Crystallography, 1974, Vol. IV, Kynoch Press, Birmingham, England.
- (14) a. Potasek, M.J.; Debrunner, P.G.; Morrison, W.H., Jr.; Hendrickson, D.N. J. Chem. Phys., 1974, 70, 2203-2206.
 - b. Datta, S.N.; Ewig, C.S.; Van Wazer, J.R.J. Mol. Struct., 1978, 48, 407-416.
- (15) Cromer, D. T.; Liberman, D. J. Chem. Phys., 1970, 53, 1891-1898.
- (16) S.J. LaPlaca, private communication
- (17) Kasper, J.S.; Interrante, L.V.; Secaur, C.A. J. <u>Amer. Chem. Soc Soc.</u>, 1975, 97, 890-891.
- (18) a. Brown, R.D.; Nunn, E.K. Austral. J. Chem., 1966, 19, 1567-1576.
 - b. Migchelsen, T.; Vos, A. Acta. Cryst., 1967, 22, 812-815.
 - c. Rundle, R.E. Acta Cryst., 1961, 14, 585-589.
- (19) a. Torrance, J.B.; Silverman, B.D. Phys. Rev. B, 1977, 15 788-801.
 - b. Silverman, B.D. Phys. Rev. B., 1978, 17, 2482-2485.
 - c. Epstein, A.J.; Lipari, N.O.; Sandman, D.J.; Nielsen, P. Phys. Rev. B, 1976, 13, 1569-1579.
 - d. Metzger, R.M. in reference 3c, p. 145-164, and references therein.
- (20) a. Silverman, B.D. J. Chem. Phys., 1979, 70, 1614-1620.
 - b. Salahub, D.R.; Messmer, R.P.; Herman, F. Phys. Rev. B, 1976, 13, 4252-4257.
- (21) Interrante, L.V.; Browall, K.W.; Hart, H.R., Jr.; Jacobs, I.S.; Watkins, G.D.; Wee, S.H. J. Amer. Chem. Soc., 1975, 97, 889-890.

Minn

Table 1 Fractional Atomic Coordinates and Anisotropic Thermal Parameters

atom	×	y	8	β12	β22	β33	β12	P ₁₃	f.23
(C)	. 34364(5)	- 12731(2)	. 62281(6)	. 01029(6)	. 00230(1)	.01948(10)	.00085(2)	.00253(6)	00117(3)
1(2)	36909(4)	. 02401(2)	.70257(5)	.00699(5)	,00205(1)	,01434(8)	.00016(2)	.00241(5)	. 00072(2)
1(3)	.37473(5)	. 17644(2)	,79679(7)	. 01207(2)	.00180(1)	. 02430(11)	00078(2)	.00441(7)	.00130(3)
S (1)	2568(2)	.0726(8)	,1492(2)	. 0077(2)	.00171(4)	. 0188(3)	00062(7)	.0035(2)	00033(9)
S(2)	.0096(2)	. 0611(8)	.3138(2)	. 0095(2)	.00188(4)	.0130(3)	.00075(7)	.0036(2)	00029(8)
S (3)	-,0098(2)	-,1074(8)	.2256(2)	. 0088(2)	.00178(4)	. 0148(3)	-, 00053(7)	. 0032(2)	00029(8)
S(4)	2390(2)	0942(8)	.0646(2)	,0082(2)	. 00175(4)	. 0161(3)	. 00062(7)	.0034(2)	.00036(8)
C (3)	. 1204(6)	-,0530(3)	,1738(7)	(9)0900	. 00162(14)	(8)0600'	. 0005(2)	0013(6)	00070(8)
C (2)	,1306(6)	0188(3)	.2127(7)	. 0056(6)	. 00166(14)	(6)0600′	.0001(2)	,0007(6)	.0000(3)
C (3)	2024(8)	1492(3)	. 2370(9)	. 0123(10)	. 00149(15)	(81)6610	0008(3)	.0010(14)	0001(3)
C(4)	.0905(8)	,1434(3)	.3132(9)	. 0157(11)	. 00157(16)	.0161(12)	. 0007(4)	, 0024(10)	0009(4)
C(5)	0529(8)	1848(3)	.1436(9)	0108(9)	.00154(15)	, 0198(14)	~ 0005(3)	, 0005(10)	. 0010(4)
C (6)	,1653(8)	-,1783(3)	. 0701(9)	. 0123(9)	.00186(17)	. 0177(13)	.0007(3)	. 0014(10)	0001(4)
H(3)	. 255(10)	,193(4)	.209(11)						
H(4)	(01)680′	.186(4)	.356(11)						
H(5)	-,006(10)	230(4)	. 162(11)						
H(6)	.218(10)	218(4)	. 013(11)						

Table II. Selected Distances and Angles in $\ensuremath{\mathsf{TTF}} \cdot I_3$

Bond Distances (1)

I(1) - I(2)	2.907(1)	S(3) - C(5)	1. 743(7)
I(2) - I(3)	2.953(1)	S(4) - C(1)	1. 731(6)
S(1) - C(2)	1. 715(6)	S(4) - C(6)	1. 730(7)
S(1) - C(3)	1. 718(7)	C(1) - C(2)	1.382(7)
S(2) - C(2)	1. 718(6)	C(3) - C(4)	1.326(11)
S(2) - C(4)	1. 724(7)	C(5) - C(6)	1. 317(11)
S(3) - C(1)	1. 713(6)		
	Non-Bonded Con	tacts (Å)	
I(1) - S(2)'	3.686(2)	I(3) - S(4)	3.870(2)
- S(4)""	3.832(2)	I(1) - C(1)	3.898(5)
- S(1)''	3.911(2)	I(2) - C(2)	3.955(5)
I(2) - S(3)'	3.881(2)	I(3) - C(4)	3.933(7)
- S(2)'	3.882(2)	I(3) - C(6)	3.902(7)
- S(1)"	3.985(2)	S(1) - S(3) ""	3.351(2)
- S(4)"	3.946(2)	S(2) - S(4)""	3.372(2)
- S(1)"	3.959(2)	$C(1) - C(2)^{M/}$	3.434(7)
-S(4)**	3.989(2)	C(3) - C(5)""	3.424(9)
I(3) - S(3)' - S(1)"	3.637(2) 3.732(2)	C(4) - C(6)""	3.439(10)
	Bond Angles		
I-I-I	175.39(2)	S(3) - C(1) - S(4)	115.1(2)
S(1) - C(2) - C(1)	122.1(4)	S(1) - C(2) - S(2)	115.7(3)
S(2) - C(2) - C(1)	122.1(4)	C(4) - C(3) - S(1)	116.6(5)
S(3) - C(1) - C(2)	123.3(4)	C(3) - C(4) - S(2)	118.1(5)
S(4) - C(1) - C(2)	121.5(4)	C(6) - C(5) - S(3)	• •
C(2) - S(1) - C(3)	95.2(3)	C(5) - C(6) - S(4)	117.4(5)
C(2) - S(2) - C(4)	94.2(3)		
C(1) - S(3) - C(5)	95.2(3)		
C(6) - S(4) - C(1)	95.1(3)		

··· MENTER.

Table III. Dimensions of TTF molecule averaged over assumed mmm molecular symmetry.

TTFBr	1. 393 Å 1. 720 1. 724 1. 332	122.3° 115.5 95.1 117.2	
TTF. L ₃ d 295° K	1. 382(7)Å 1. 719(8) 1. 728(11) 1. 322(11)	122. 5(8)° 115. 4(4) 94. 9(5) 117. 3(6)	
TTF.L.	1.350Å 1.732 1.721 1.336	122.9 114.3 95.7 117.2	ω γ ω
TTF-TCNQ ^b 298° K	1. 369(4) Å 1. 743(4) 1. 736(5) 1. 323(4)	122. 6(3) 114. 7(3) 94. 9(3) 117. 6(4)	C a C P
TTF ^a 295°K	1. 349(3) 1. 757(2) 1. 726(4) 1. 314(3)	122.8(2) 114.5(2) 94.4(2) 118.3(4)	O=O
	פיט ב א	0 1 2 B	

a Phillips, T.E.; Kistenmacher, T.J.; Ferraris, Cowan, D.O. Chem. Comm. 1973, 14, 471-472.

b Kistenmacher, T.J.; Phillips, T.E.; Cowan, D.O. Acta. Crystallogr, 1974, B30, 763-768.

c Obtained by Fourier refinement and comparable standard errors are unavailable. Ref. 6c.

d This work

e Reference 16.

Table IV. Bond Angles and Distances in Triiodide Ions

$$I \xrightarrow{a} I \xrightarrow{b} I$$

	⊕ (deg)	a(Å)	b(Å)
TTF·I ₃ a	175.39(2)	2.9533(8)	2.9078(7)
$((C_6H_5)CONH_2)_2H^+I_3^-$	177. 2(5)	2, 959(8)	2.900(8)
•	176.5(4)	2.943(8)	2.921(8)
$(C_6H_5)_4AsI_3$	175.61(5)	2.920(2)	2.920(2)
CsI ₃ e	177. 9(1)	3.042(4)	2.840(4)
NH ₄ I ₃ ^d	180. 0(1)	3.113(4)	2.791(4)
(thiocarbamato) ₂ CuI ₃	176.5(3)	2.919(7)	2.899(7)
$(phenacetin)_2 \cdot H^+ I_3^- \cdot I_2^f$	180	2.907(1)	2.907(1)

This work.

b This structure contains parallel chains of triiodide ions, with two types of I₃ moeties; Reddy, J.M.; Knox, K.; Robin, M.B. J. Chem. Phys., 1964, 40, 1082-1089.

Runsink, J.; Swen-Walstra, S.; Migchelsen, T. Acta. Crystaliogr., Sect. B, 1972, 28, 1331-1335.

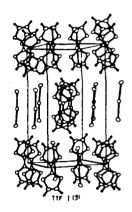
Cheesman, G.H.; Finney, A.J.T., <u>Acta. Crystallogr.</u>, <u>Sect. B</u>, <u>1970</u>, 26, 904-906.

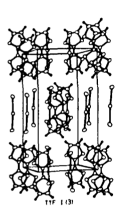
Wijnhoven, J.G.; Van den Hark, T.E.M.; Beurkeus, P.T. J. Cryst. Mol. Struct., 1972, 2, 189-196.

Herbstein, F. H.; Kapon, M. Nature (London), 1972, 239, 153-1954.

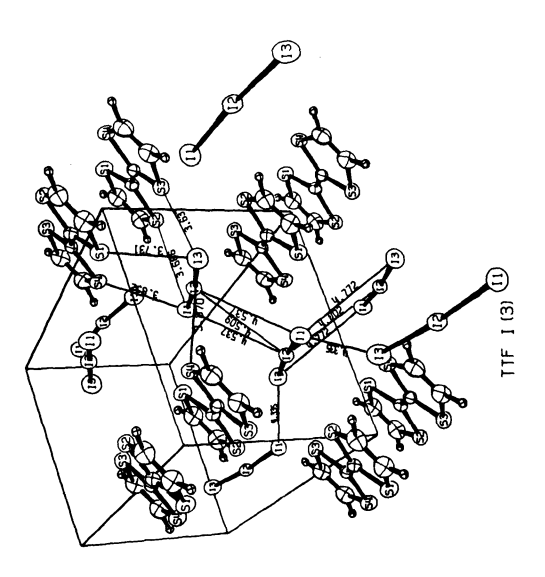
FIGURE CAPTIONS

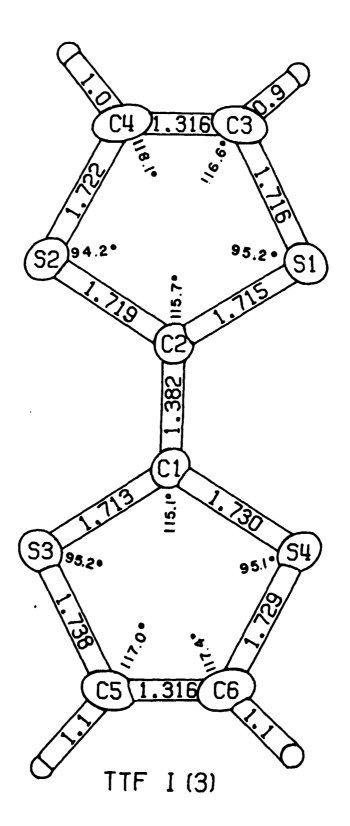
- Figure 1. Stereoscopic view of the molecular packing of $TTF \cdot I_3$. The monoclinic b axis is vertical.
- Figure 2. View of stacking within a sheet of $(TTF)_2^{+2}$ dimers and triiodide ions in the ac plane. The zig-zagging triiodide chains along b also are shown.
- Figure 3. Bond distances and angles in the TTF⁺ ion. Probability ellipsoids (50%) and the numbering scheme are also shown.





a describe





Paralle Market

- Kenda

TECHNICAL REPORT DISTRIBUTION LIST, GEN

	<u>No</u> . Copies		<u>No</u> . Copie:
	ODDIES		<u>oopto</u>
Office of Naval Research		U.S. Army Research Office	
Attn: Code 472		Attn. CRD-AA-IP	
800 North Quincy Street		P.O. Box 1211	
Arlington, Virginia 22217	2	Research Triangle Park, N.C. 27709	1
ONR Branch Office		Naval Ocean Systems Center	
Attn: Dr. George Sandoz		Attn: Mr. Joe McCartney	
536 S. Clark Street		San Diego, California 92152	1
Chicago, Illinois 60605	1		
OND TO A COCCUM		Naval Weapons Center	
ONR Branch Office		Attn: Dr. A. B. Amster,	
Attn: Scientific Dept.		Chemistry Division	
715 Broadway	•	China Lake, California 93555	1
New York, New York 10003	1	3 1 G/ /1 B / 1 1 1 1 1	
		Naval Civil Engineering Laboratory	
ONR Branch Office		Attn: Dr. R. W. Drisko	4
1030 East Green Street	•	Port Hueneme, California 93401	1
Pasadena, California 91106	1		
		Department of Physics & Chemistry	
ONR Branch Office		Naval Postgraduate School	•
Attn: Dr. L. H. Peebles		Monterey, California 93940	1
Building 114, Section D		D 4 1 01 - 611	
666 Summer Street	•	Dr. A. L. Slafkosky	
Boston, Massachusetts 02210	1	Scientific Advisor	
M		Commandant of the Marine Corps	
Director, Naval Research Laboratory		(Code RD-1)	1
Attn: Code 6100	•	Washington, D.C. 20380	1
Washington, D.C. 20390	1	Office of Naval Research	
The Analogous Company		Attn: Dr. Richard S. Miller	
The Assistant Secretary		800 N. Quincy Street	
of the Navy (R,E&S)		Arlington, Virginia 22217	1
Department of the Navy		ALLINGTON, VIIGINIA 22217	•
Room 4E736, Pentagon	1	Naval Ship Research and Development	
Washington, D.C. 20350	4	Center	
Commander, Naval Air Systems Command		Attn: Dr. G. Sosmajian, Applied	
Attn: Code 310C (H. Rosenwasser)		Chemistry Division	
Department of the Navy		Annapolis, Maryland 21401	1
Washington, D.C. 20360	1	, ,	
"Estimated by the costs	-	Naval Ocean Systems Center	
Defense Documentation Center		Attn: Dr. S. Yamamoto, Marine	
Building 5, Cameron Station		Sciences Division	
Alexandria, Virginia 22314	12	San Diego, California 91232	1
		•	
Dr. Fred Saalfeld		Mr. John Boyle	
Chemistry Division		Materials Branch	
Maval Research Laboratory		Naval Ship Engineering Center	
Washington, D.C. 20375	1	Philadelphia, Pennsylvania 19112	1

A Printer

TECHNICAL REPORT DISTRIBUTION LIST, GEN

No. Copies

Dr. Rudolph J. Marcus
Office of Naval Research
Scientific Liaison Group
American Embassy
APO San Francisco 96503

Mr. James Kelley
DTNSRDC Code 2803
Annapolis, Maryland 21402

ALICE AND A

TECHNICAL REPORT DISTRIBUTION LIST, 053

	No. Copies		No. Copies
Dr. R. N. Grimes University of Virginia Department of Chemistry Charlottesville, Virginia 22901	1	Dr. M. H. Chisholm Department of Chemistry Indiana University Bloomington, Indiana 47401	1
Dr. M. Tsutsui Texas A&M University		Dr. B. Foxman Brandeis University	
Department of Chemistry College Station, Texas 77843	1	Department of Chemistry Waltham, Massachusetts 02154	1
Dr. M. F. Hawthorne University of California Department of Chemistry Los Angeles, California 90024	1	Dr. T. Marks Northwestern University Department of Chemistry	
Dr. D. B. Brown		Evanstor, Inlinois 60201	1
University of Vermont Department of Chemistry Burlington, Vermont 05401	1	Dr. G. Geoffrey Pennsylvania State University Department of Chemistry	
or. W. B. Fox Naval Research Laboratory		University Park, Pennsylvania 16802	1
Chemistry Division Code 6130 Washington, D.C. 20375	1	Dr. J. Zuckerman University of Oklahoma Department of Chemistry Norman, Oklahoma 73019	1
Dr. J. Adcock	•	Professor 0. T. Beachley	•
University of Tennessee Department of Chemistry		Department of Chemistry State University of New York	
Knoxville, Tennessee 37916	1	Buffalo, New York 14214	1
Dr. A. Cowley University of Texas Department of Chemistry		Professor P. S. Skell Department of Chemistry	
Austin, Texas 78712	1	The Pennsylvania State University University Park, Pennsylvania 16802	1
Dr. W. Hatfield University of North Carolina Department of Chemistry		Professor K. M. Nicholas Department of Chemistry	
Chapel Hill, North Carolina 27514	1	Boston College Chestnut Hill, Massachusetts 02167	1
Dr. D. Seyferth Massachusetts Institute of Technology		Professor R. Neilson Department of Chemistry	
Department of Chemistry Cambridge, Massachusetts 02139	1	Texas Christian University Fort Worth, Texas 76129	1
Professor H. Abrahamson University of Oklahoma		Professor M. Newcomb Texas A&M University Department of Chemistry	
Department of Chemistry Norman, Oklahoma 73019	1	Department of Chemistry College Station, Texas 77843	1